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(54) Liquid fabric conditioner and dryer sheet fabric conditioner containing compatible silicones.

(57) Liquid fabric conditioning compositions and fabric conditioning composition for coating a flexible substrate for subsequent use in a mechanical tumble dryer are disclosed. The compositions incorporate organosilicones which form compatible mixtures with common fabric softening agents.

BACKGROUND OF THE INVENTION

FIELD OF THE INVENTION

A first aspect of the instant invention relates to conditioning of fabrics in an aqueous wash bath, to liquid compositions containing fabric conditioning ingredients and to processes for making the compositions. A second aspect of the invention relates to application of adjuvants to fabrics in tumble-dryer automatic dryers. More particularly, to an article in the form of a flexible substrate carrying a fabric conditioning composition.

RELATED ART

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Silicones have been applied to fabrics during manufacture of fabrics or during the make up of articles of clothing. With respect to application of silicones to fabrics during a laundry process, Great Britain Patent Application 1,549,480; Burmeister et al., U.S. Patent 4,818,242; Konig et al., U.S. Patent 4,764,089; Konig et al., U.S. Patent 4,806,255; Dekker et al., U.S. Patent 4,661,267 and Trinh et al., U.S. Patent 4,661,269 describe aqueous dispersions or emulsions of certain silicones of limited viscosity incorporated in liquid rinse-cycle fabric softening compositions. A fabric softening composition containing emulsified silicone is also taught by Barrat et al. in U.S. Patent 4,446,033. Coffindafer et al., U.S. Patent 4,800,026 discloses fabric care compositions containing curable amine functional silicones.

The application of fabric softeners to fabrics in the tumble dryer by use of a flexible substrate carrying the fabric softeners is known in the art. The advantages of dryer added fabric conditioning include a more convenient time of addition in the laundry process and avoidance of undesirable interaction of softening agents with detergents.

Rudy et al., U.S. Patent 3,972, 131 discloses dryer sheets including a silicone oil as an ironing aid. Kasprzak et al., U.S. Patent 4,767,548 discloses the use of certain silicones in dryer sheet formulations. Coffindafer et al., U.S. Patent 4,800,026 discloses curable amine functional silicones in fabric care compositions.

In the manufacture of the dryer added fabric conditioning sheets described in the references mentioned above, when silicones are mixed with fabric softeners, the resulting mixtures are non-homogeneous and phase separation occurs readily. The homogeneity of such mixtures is ensured only by continuous vigorous agitation. An additional problem associated with the use of a nonhomogeneous mixture is the separation of actives at the point of application of the active mixture on the substrate resulting in unevenly impregnated sheets.

The compositions disclosed in the art contain individual particles of a silicone and individual particles of a fabric softening agent.

In the present invention the dispersed particle is a composite particle containing a mutually compatible mixture of a silicone and a fabric softening component. Compatible organosilicones described herein preferably form mutually soluble mixtures with certain types of commonly used fabric softening agents. Critically, the organosilicones in the dispersed composite particles do not separate from fabric softening agents during processing, on standing, during coating or drying of the dryer sheet. An additional advantage afforded by the present invention is a simplified manufacture of fabric softener compositions since the silicones need not be dispersed separately and can be introduced into the composition simultaneously with a fabric softener.

Another advantage of using compatible silicones is that compatible silicones enhance the spreading of the fabric softening agents on the fabric surface as compared to the spreading of the fabric softening agents alone or in combination with incompatible silicones. As a result of the use of compatible silicones as described herein greater, more complete surface coverage by a fabric softening agent is achieved with a further advantage of smaller dosage requirements. Additionally, even and uniform distribution of the actives on the dryer sheet can be attained, alleviating the problem of unevenly impregnated sheets.

Accordingly, it is an object of the present invention to provide a liquid fabric conditioning composition which contains composite particles of a mutually soluble mixture of a fabric softening component and an organositicone.

It is a further object of the invention to provide processes by which the aforementioned composition can be manufactured.

It is a further object of the Invention to provide an article which provides for release of a fabric conditioning composition within an automatic laundry dryer, the composition containing a compatible mixture of a fabric softening component and a selected organosilicone.

These and other objects and advantages will appear as the description proceeds.

SUMMARY OF THE INVENTION

The present invention is based, in part, on the discovery that specific silicones, defined herein as compatible, are capable of forming compatible mixtures with certain conventional fabric softening agents.

It is important to differentiate between compatible and incompatible silicones and between mutually soluble and insoluble mixtures of silicones and fabric softeners. Compability as taught herein is critical and is ascertained by the appearance of the mixture of a silicone and a fabric softener. When a silicone and a fabric softener are heated and mixed together, the resulting liquid mixtures are either transparent or opaque. In the transparent mixtures, silicone and fabric softener are mutually soluble and are, accordingly suitable for use in the present invention. In the opaque mixtures, silicone and fabric softener are mutually insoluble and the mixtures may or may not form mutually stable dispersions. A mutually stable dispersion is also compatible and is formed if a mixture of a silicone and a fabric softener does not separate into more than one phase on storage at elevated temperatures and if the mixture does not form a uniform solld or liquid on cooling. Thus, the class of compatible mixtures as defined herein includes mutually soluble mixtures as well as mutually stable dispersions. Compatibility of the mixture is critical and is determined by the silicone softener compatibility test (SSCT) described below.

In its broadest aspect, some objects of the invention are accomplished by a liquid fabric conditioning composition which includes about 1% to about 60% of composite particles containing a mutually soluble mixture of a fabric softening component and an organosilicone. Of course, these particles can also be added to a liquid containing other fabric treating ingredients, including for example, softeners.

Other objects of the invention are accomplished by an article comprising a flexible substrate carrying an effective amount of a fabric conditioning composition affixed thereto in a manner which provides for release of the conditioning composition within an automatic tumble dryer at dryer operating temperatures.

The fabric softening component employed herein for liquid compositions may be any commonly used fabric softening agent complying with the above conditions provided that it must include at least a portion of cationic quaternary ammonium salts either used singly or, optionally, in admixture with other softening agents such as nonionic softeners selected from the group of tertiary amines having at least one C₈₋₃₀ alkyl chain, esters of polyhydric alcohols, fatty alcohols, ethoxylated fatty alcohols, alkyl phenols, ethoxylated alkylphenols, ethoxylated fatty amines, ethoxylated monoglycerides, ethoxylated diglycerides, mineral oils, polyols, carboxylic acids having at least 8 carbon atoms, and mixtures thereof.

The fabric conditioning composition employed in articles of the present invention contains (A) certain fabric softening agents used singly or in admixture with each other and (B) an organositicone having specific structural requirements and a specific %CH₂ content.

DETAILED DESCRIPTION OF THE INVENTION

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The fabric conditioning compositio of the present invention includes a cationic quaternary ammonium salt. The couterion is methyl sulfate or any halide.

Examples of cationic quaternary ammonium salts include, but are not limited to:

- 1. Acyclic quaternary ammonium salts having at least two C_8 to C_{90} , preferably C_{12} to C_{22} alkyl chains, such as: ditallow/dimethyl ammonium chloride, di(hydrogenated tallow)dimethyl ammonium chloride, distearyl-dimethyl ammonium chloride, discocodimethyl ammonium chloride and the like;
- Cyclic quaternary ammonium salts of the imidazolinium type such as di(hydrogenated tallow)dimethyl imidazolinium methyl sulfate, 1-ethylene-bis(2-tallow-1-methyl) imidazolinium methyl sulfate and the like;
- 3. Diamido quaternary ammonium salts such as:
- methyl-bis(hydrogenated tallow amidoethyl)-2-hydroethyl ammonium methyl sulfate,
- methyl-bis(tallowamidoethyl)-2-hydroxypropyl ammonium methyl sulfate and the like;
- 4. Blodegradable quaternary ammonium salts such as N,N-di(tallowoyl-oxy-ethyl)-N,N,-dimethyl ammonium chloride, and N,N-di(tallowoyl-oxy-propyl)-N,N-dimethyl ammonium chloride and the like. When fabric conditioning compositions employ biodegradable quaternary ammonium salts, the pH of the composition is preferably adjusted to between about 2 and about 5. Biodegradable quaternary ammonium salts are described, for example, in U.S. Patents 4,767,547 and 4,789,491 incorporated by reference herein.
- 5. Mixtures of water-insoluble cationic fabric softeners and polyalkoxylated ammonium salts as described in U.S. Patent 4,422,949 incorporated by reference herein. Such mixtures are particularly suitable for incorporation in concentrated form of the liquid compositions herein.

The fabric softening component may include other fabric softeners in addition to the cationic quaternary

ammonium salts. Additional fabric softeners suitable for use herein can be selected from the following classes of compounds:

 Tertiary fatty amines having at least one and preferably two C₈ to C₃₀, preferably C₁₂ to C₂₂ alkyl chains. Examples include hardened tallow amine and cyclic amines such as 1-(hydrogenated tallow)amidoethyl-2-(hydrogenated tallow) imidazoline. Cyclic amines which may be employed for the compositions herein are described in U.S. Patent 4,806,255 incorporated by reference herein.

ii. Carboxylic acids having 8 to 30 carbon atoms and one carboxylic group per molecule. The alkyl portion has 8 to 30, preferably 12 to 22 carbon atoms. The alkyl portion may be linear or branched, saturated or unsaturated, with linear saturated alkyl preferred. Stearic and myristic acids are preferred fatty acids for use in the composition herein. Examples of these carboxylic acids are commercial grades of stearic acid and the like which may contain small amounts of other acids.

iii. Esters of polyhydric alcohols such as sorbitan esters or glycerol stearate. Sorbitan esters are the condensation products of sorbitol or iso-sorbitol with fatty acids such as stearic acid. Preferred sorbitanesters are monoalkyl. A common example of sorbitan ester is SPAN 60 (ICI) which is a mixture of sorbitan and isosorbide stearates.

iv. Fatty alcohols, ethoxylated fatty alcohols, alkylphenols, ethoxylated alkyl phenols, ethoxylated fatty amines, ethoxylated monoglycerides and ethoxylated digiycerides.

v. Mineral oils, and polyols such as polyethylene glycol.

vi. Condensation products of higher fatty acids with polyamines, selected from the group consisting of hydroxyalkyl alkylene diamines, dialkylene triamines and mixtures thereof, as described in U.S. Patent 4,661,269, the disclosure of which incorporated by reference herein.

Preferred fabric softeners for use herein are acyclic quaternary ammonium salts, ditallowdimethyl ammonium chloride being most preferred for fabric conditioning compositions of this invention. Especially preferred are mixtures of ditallowdimethyl ammonium chloride with fatty acids, particularly stearic acid or myristic acid.

About 1% to about 40% of the fabric softening component is used in the compositions of the invention. There must be included at least a sufficient amount of quaternary ammonium salt to achieve anti-static effect, for example, about 1% to 3% in the dilute product and about 2% to about 5% in the concentrated product. On the other hand, the entire fabric softening component may be quaternary ammonium salt. The diluted version of the product contains about 1% to about 12%, preferably about 3% to about 10% and most preferably about 4% to about 7% of the fabric softening component. The concentrated version of the product contains about 13% to about 40%, preferably about 13% to 30% and most preferably about 13% to about 20% of the fabric softening component.

When the fabric softening composition is carried on a flexible substrate according to the article of the present invention the fabric softening agents (A) include conventionally used cationic and nonionic fabric softening agents such as those listed in (1) to (5) and (i) to (vi) above.

The amount of the fabric softening composition on the sheet is subject to normal coating parameters such as, for example, viscosity and melting point of the fabric softening components and is typically about 0.5 grams to about 5 grams, preferably about 1 gram to about 3.5 grams. The fabric softening composition employed in the present invention contains about 0.1% to about 95% of the fabric softening component. Preferably from about 10% to about 80% and most preferably from about 30% to about 70% of the fabric softening component is employed herein to obtain optimum softening at minimum cost. When the fabric softening component includes a quaternary ammonium salt, the salt is used in the amount of about 10% to about 80%, preferably about 30% to about 70%.

Silicone

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The second essential ingredient of the fabric softening composition employed in the present invention is a compatible organosilicone.

The organosilicones employed in the present invention (also termed herein as compatible silicones) are capable of forming compatible mixtures with the fabric softeners listed above.

The organosilicones employed herein have a %CH₂ content of about 25% to about 90%. The % CH₂ content is defined as

 $%CH_2 = \frac{\text{number of methylene}(CH_2) \text{ groups}}{\text{number of methylene groups and methyl groups}} \times 100\%$

The organosilicones included in the fabric conditioning compositions of the invention contain at least one unit of Formula A:

wherein m is a number from 0 to 2 and R is a mono valent hydrocarbon radical.

The value of (3-m)/2 in Formula A means the ratio of oxygen atoms to silicon atoms, i.e. SiO_{1/2} means one oxygen is shared between two silicon atoms.

R1 in Formula A is selected from the group consisting of:

i. a hydrocarbon radical having from 6 to 45 carbon atoms, preferably from 8 to 18 carbon atoms and which may be saturated, unsaturated, cyclic, acyclic, alkyl or aromatic:

ii. a unit of Formula A1

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$$-(CH_2)_a - (N-CH_2CH_2)_b - N-R^2$$

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wherein a is a number of at least 1, preferably 3; b is a number from 0 to 10, preferably 1; R2 is

R³ is a hydrocarbon radical having from 4 to 40 carbon atoms preferably from 8 to 18 carbon atoms and may be saturated, unsaturated, cyclic, acyclic, alkyl or aromatic; and R⁴ is hydrogen or a hydrocarbon radical having from 1 to 40 carbon atoms, preferably hydrogen; and iii. a unit of Formula A2

wherein R^6 and R^6 are independently selected from hydrogen or a hydrocarbon radical having from 1 to 45 carbon atoms which may be saturated, unsaturated, cyclic, acyclic, alkyl or aromatic and at least one of R^6 and R^6 is a hydrocarbon radical having from 6 to 45 carbon atoms, R^7 is

wherein R⁸ is a divalent organic radical having from 1 to 12 carbon atoms which may be saturated, unsaturated, cyclic, acyclic, alkyl or aromatic, and preferably is -CH₂CH₂CH₂-O-CH₂.

Thus, organosilicones employed in the present invention include alkylsilicones and alkylaminosilicones which satisfy the structural parameters described above and which have % methylene (%CH₂) content of about 25% to about 90%. Compatibility of the organosilicones herein with fabric softening agents depends, in part, on the %CH₂ content of the organosilicones. The preferred range of the %CH₂ content for the silicones herein is from about 40% to about 90%, more preferably from about 50% to about 85%, and most preferably from about 50% to about 75% to increase the degree of compatibility of the mixtures containing relatively large amounts of silicone.

The organosilicones included in the compositions herein may be linear, branched, or partially crosslinked, preferably linear, and may range from fluid, liquid or viscous liquid, gum and solid.

An example of an alkylsilicone suitable for use herein is:

An example of a suitable alkylaminositicone containing the unit of Formula A1 is:

An example of an alkylaminosilicone containing the unit of Formula A2 is:

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Alkylsilicones employed in this invention may be produced by reacting a hydrosiloxane co-polymer with a hydrocarbon having 6 to 45 carbon atoms and having a terminal vinyl functionality. Such reactions are described, for example, in Chemistry and Technology of Silicones by Walter Notl, Academic Press, N.Y. (1968), pages 49-51 and 219-226. Commercially available alkylsilicones suitable for use herein are, for example, Masil 264, Masil 265 HV from Mazer International Corp. or ABIL - Wax 9800/or ABIL - Wax 9801 from Th. Goldschmidt AG.

Alkylaminosilicones employed in this invention may be produced by 1) treating silicones containing primary or secondary amine functional groups with epoxides such as ethylene oxide to form alkylaminosilicones having the unit of Formula A1, or 2) by treating epoxysilicones with primary or secondary amines such as dioccoamine to form alkylaminosilicones having the unit of Formula A2.

The modified alkylaminosilicones of the invention having the unit of Formula A1 may be prepared by mixing epoxide compounds with aminosilicones in a pressure reactor and heating for about 24 hours, after which the unreacted epoxide compound is vacuum stripped off. The amount of epoxide to be used is calculated based upon the number of amine functional groups on the alkylaminosilicone. Preferably, two epoxides are reacted for every primary amine and one epoxide for every secondary amine, in order to convert them to tertiary amines. A stoichiometric amount or up to 25% excess if epoxide can be used. The reaction is preferably conducted between 25°C and 150°C, especially between 50°C and 100°C. The pressure is preferably maintained from 50 psi to 300 psi, particularly from 50 psi to 150 psi. Typical aminosilicone starting compounds would include Dow Corning Q2-8075. The art of making alkylaminosilicones having the unit of Formula A1 is disclosed in Examples 1 and 2 herein and in the copending patent application of Line 1al. entitled "Hydroxylhydrocarbyl Modified Aminoalkyl Silicones', S.N. 449,360 filled December 6, 1989.

The modified alkylaminosilicones having the unit of Formula A2 may be prepared by mixing epoxysilicones, secondary amines, and a solvent such as isopropanol or toluene, and heating the mixture at reflux for about 24 hours, after which the solvent is removed by distillation or vacuum stripping. The amount of amine to be used is calculated based upon the number of epoxy functional groups on the epoxysilicone. Preferably, one secondary amine is reacted for every epoxy functional group in order to convert the amine to tertiary amine. A stoichiometric amount or up to 25% excess of amine can be used. The reaction is preferably conducted between 50°C and 150°C, especially between 75°C and 110°C. The reaction is preferably conducted at atmospheric pressure, but may be conducted in a pressure reactor with the pressure being maintained from 50 psi to 300 psi.

The modified alkylaminosilicones employed in this invention contain amine groups which may be quaternised with, for example, alkyl halide or methyl sulfate, or may be protonated with a Lewis acid such as hydrochloric acid, acetic acid, citric acid, formic acid and the like. Alkylsilicones and alkylaminosilicones employed herein may, in addition to the units of Formula A, contain secondary units selected from the group consisting of a unit of Formula B1 and a unit of Formula B2:

$$R^{11} - O(CH_2CHO)_CR^{10} - (SiO)_{3y/2}$$
 Formula B1

$$R^{11} - O(CH_2CH_2CH_2CHO)_d - R^{10} - (SiO)_{3z/2}$$
 Formula B2

wherein R¹¹ is a hydrocarbon radical having from 1 to 40 carbon atoms, preferably is CH₃; R⁹ is a hydrocarbon radical having from 1 to 3 carbon atoms; R¹⁰ is oxygen or alkylene having from 1 to 8 carbon atoms, preferably propylene; c and d are numbers from 0 to 50, preferably 2 to 15; and y and z are numbers from 0 to 2.

Organosilicones preferred for use herein have a %CH₂ content of about 40% to about 90% and are either alkylaminosilicones having the unit of Formula A1 or alkylsilicones.

The amount of organosilicone employed herein generally ranges from about 0.1% to about 20%, and is preferably at least about 0.5% to about 2% to maximise the spreading of the fabric softeners on fabric surface, but could be higher in concentrated liquids. Preferably when the organosilicone is carried on a substrate, the amount employed is from 0.1% to 20%, more preferably at least 3% to 20%. The amount of the organosilicone is governed by the ratio at which the mutually soluble mixture of the fabric softening component and the organosilicone is formed.

The weight ratio of the organosilicone to the fabric softening component in the fabric conditioning compositions employed herein is from about 100:2 to about 1:100, preferably from about 2:100 to about 20:100, but must be such that a compatible mixture can be formed.

Silicone/Softener Compatibility Test (SSCT)

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As described above, mixtures defined as compatible herein include mutually soluble as well as mutually stable dispersible mixtures. Compatibility of the fabric conditioning mixtures herein depends on the structure and the %CH₂ content of the organosilicone and the particular fabric softeners employed in the mixture. SSCT provides a basis for selecting appropriate combinations of the fabric softening component and the organosilicone.

The test may be used to determine the compatibility at a particular weight ratio of interest or to determine a minimum concentration of the silicone at which a compatible mixture of the silicone and the fabric softening component is formed.

SSCT is conducted as follows:

a 10 gram sample of the fabric softener or a combination of fabric softeners is placed into a clear glass flask equipped with a stirring mechanism, such as a magnetic stirrer. If either the fabric softener or the silicone is a solid at room temperature, it is melted before the test is begun with the test taking place above the melting point of the fabric softener or the silicone. The silicone of interest is slowly introduced with, conveniently, a Pasteur pipet into the flask, with stirring. It is estimated that the weight of one drop represents about 1% silicone concentration, so the silicone is mixed with the fabric softener 1% at a time. Thus, the lowest concentration of the silicone in the mixture is about 1%.

If the resulting mixture of the fabric softening agent and the silicone stays clear over the entire investigated range of the silicone, this indicates that the components of the mixture are mutually soluble over the investigated concentration range and, accordingly, are compatible. Clear mixtures are defined herein as mixtures having about 90% transmittance when measured with a visible light probe (one centimeter pathlength) against distilled water background using Brinkman PC800 colometer.

The mixture may also become cloudy indicating that the silicone and the fabric softener are not mutually soluble at that weight % of the silicone. In this case, if the mixture became cloudy, the weight percent of the silicone added to produce cloudiness is calculated. This number, termed compatibility a, then represents the weight percent of the silicone to produce a cloudy mixture. Cloudy samples are placed in an oven at 100°C for at least two hours, then cooled to room temperature and inspected. Samples which have completely separated

into distinct layers are incompatible and are not useful for the invention. Samples which maintain a stable, dispersed character are compatible and, hence, useful in the invention.

It is sufficient, for practical applications, to investigate the silicone concentration range of up to about 30%. However, the entire range up to 100% of the silicone concentration may be investigated if desired. When the entire range of the silicone concentration is to be investigated, the silicone is added until the mixture contains about 60% by weight of the silicone. Silicone addition is then stopped, and the experiment is repeated by adding the fabric softener to a 10 gram sample of the silicone. In those samples that became cloudy, the weight percent of the softener added to produce cloudiness is calculated and subtracted from 100, the resulting number is termed herein compatibility 6.

 α compatibility reflects compatibility of the mixtures containing a fabric softener as a major component, whereas β compatibility reflects compatibility of the mixtures containing a silicone as a major component. Minimal difference between β and α (β - α) reflects degree of compatibility of the mixture: more compatible mixtures have a lower number for β - α .

Preferably, the silicone and the fabric softening component are compatible at a silicone concentration of at least about 2%.

Mutually soluble and clear mixtures of the silicone and the fabric softening component indicate the highest degree of compatibility and are preferred.

Various additives may be used in combination with the compatible mixture of the fabric softening component and the compatible silicone. The additives are used in the amounts that do not substantially affect the compatibility of the mixture and include small amounts of incompatible silicones, such as predominantely linear polydialkylsioxanes, e.g. polydimethylsiloxanes; soil release polymers such as block copolymers of polyethylene oxide and terephthalate; fatty amines selected from the group consisting of primary fatty amines, secondary fatty amines, tertiary fatty amines and mixtures thereof; amphoteric surfactants; smectile type inoranic clavs: anionic soaps, switterionic quaternary ammonium compounds; and nonionic surfactants.

Other optional ingredients include emulsifiers, electrolytes, optical brighteners or fluorescent agents, buffers, perfumes, colourants, germicides and bactericides.

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The fabric conditioning compositions of the invention can be used in the rinse cycle of a conventional home laundry operation. Generally, rinse water has a temperature of from about 5°C to about 70°C. The concentration of the total active ingredients is generally from about 2 ppm to about 1000 ppm, preferably from about 10 ppm to about 500 ppm, by weight of the aqueous rinsing bath. When multiple rinses are used, the fabric conditioning composition is preferably added to the final rinse.

Additionally, an article is disclosed for conditioning fabrics in a tumble dryer. The article of the invention comprises a flexible substrate which carries a fabric conditioning amount of a conditioning composition and is capable of releasing the conditioning composition at dryer operating temperatures. The conditioning composition in turn has a preferred melting (or softening) point of about 25°C to about 150°C.

The fabric conditioning composition employed in the invention is coated onto a dispensing means which effectively releases the fabric conditioning composition in a tumble dryer. Such dispensing means can be designed for single usage or for multiple uses. One such article comprises a sponge material releasably enclosing enough of the conditioning composition to effectively impart fabric softeness during several drying cycles. This multi-use article can be made by filling a porous sponge with the composition. In use, the composition melts and leaches out through the pores of the sponge to soften and condition fabrics. Such a filled sponge can be used to treat several loads of fabrics in conventional dryers, and has the advantage that it can remain in the dryer after use and is not likely to be misplaced or lost.

Another article comprises a cloth or paper bag releasably enclosing the composition and sealed with a hardened plug of the mixture. The action and heat of the dryer opens the bag and releases the composition to perform its softening.

A highly preferred article comprises the compositions containing a softener and a compatible organosilicone releasily affixed to a flexible substrate such as a sheet of paper or woven or nonwoven doth substrate. When such an article is placed in an automatic laundry dryer, the heat, moisture, distribution forces and tumbling action of the dryer removes the composition from the substrate and deposits it on the fabrics.

The sheet conformation has several advantages. For example, effective amounts of the compositions for use in conventional dryers can be easily absorbed onto and into the sheet substrate by a simple dipping or padding process. Thus, the end user need not measure the amount of the composition necessary to obtain fabric softness and other benefits. Additionally, the flat configuration of the sheet provides a large surface area which results in efficient release and distribution of the materials onto fabrics by the tumbling action of the dryer.

The substrates used in the articles can have a dense, or more preferably, open or porous structure. Examples of suitable materials which can be used as substrates herein include paper, woven cloth, and non-woven cloth. The term 'cloth' herein means a woven or non-woven substrate for the articles of manufacture, as dis-

tinguished from the term 'fabric' which encompasses the clothing fabrics being dried in an automatic dryer.

It is known that most substances are able to absorb a liquid substance to some degree; however, the term 'absorbent', as used herein, is intended to mean a substrate with a absorbent capacity (i.e., a parameter representing a substrate's ability to take up and retain a liquid) from 4 to 12, preferably 5 to 7 times its weight of water.

If the substrate is a foamed plastics material, the absorbent capacity is preferably in the range of 15 to 22, but some special foams can have an absorbent capacity in the range from 4 to 12.

Determination of absorbent capacity values is made by using the capacity testing procedures described in U.S. Federal Specification (UU-T-595b). modified as follows:

1. tap water is used instead of distilled water;

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- 2. the specimen is immersed for 30 seconds instead of 3 minutes;
- 3. draining time is 15 seconds instead of 1 minute; and
- 4. the specimen is immediately weighed on a torsion balance having a pan with turned-up edges.

Absorbent capacity values are then calculated in accordance with the formula given in said Specification. Based on this test, one-ply, dense bleached paper (e.g., Kraft or bond having a basis weight of about 32 pounds per 3,000 square feet) has an absorbent capacity of 3.5 to 4; commercially available household one-ply towel paper has a value of 5 to 6; and commercially available two-ply household towelling paper has a value of 7 to

Suitable materials which can be used as a substrate in the invention herein include, among others, sponges, paper, and woven and non-woven cloth, all having the necessary absorbency requirements defined above.

The preferred non-woven coth substrates can generally be defined as adhesively bonded fibrous or filamentous products having a web or carded fibre structure (where the fibre strength is suitable to allow carding), or comprising fibrous mats in which the fibres or filaments are distributed haphazardly or in random array (i.e. an array of fibres in a carded web wherein partial orientation of the fibres is frequently present, as well as a completely haphazard distribution orientation), or substantially aligned. The fibres or filaments can be natural (e.g. wool, silk, jute, hemp, cotton, linen, sisal, or ramie) or synthetic (e.g. rayon, celloluse ester, polyvinyl derivatives, polyolefins, polyamides, or polyesters).

The preferred absorbent properties are particularly easy to obtain with non-woven cloths and are provided merely by building up the thickness of the cloth, i.e., by superimposing a plurality of carded webs or mats to a thickness adequate to obtain the necessary absorbent properties, or by allowing a sufficient thickness of the fibres to deposit on the screen. Any diameter or denier of the fibre (generally up to about 10 denier) can be used, inasmuch as it is the free space between each fibre that makes the thickness of the cloth directly related to the absorbent capacity of the cloth, and which, further, makes the non-woven cloth especially suitable for impregnation with a composition by means of intersectional or capillary action. Thus, any thickness necessary to obtain the required absorbent capacity can be used.

When the substrate for the composition is a non-woven cloth made from fibres deposited haphazardly or in random array on the screen, the articles exhibit excellent strength in all directions and are not prone to tear or separate when used in the automatic clothes driver.

Preferably, the non-woven cloth is water-laid or air-laid and is made from cellulosic fibres, particularly from regenerated cellulose or rayon. Such non-woven cloth can be lubricated with any standard textile lubricant. Preferably, the fibres are from 5mm to 50mm in length and are from 1.5 to 5 denier. Preferably, the fibres are at least partially orientated haphazardly, and are adhesively bonded together with a hydrophobic or substantially hydrophobic binder-resin. Preferably, the cloth comprises about 70% fibre and 30% binder resin polymer by weight and has a basis weight of from about 18 to 45g per square metre.

In applying the fabric conditioning composition to the absorbent substrate, the amount impregnated into and/or coated onto the absorbent substrate is conveniently in the weight ratio range of from about 10:1 to 0.5:1 based on the ratio of total conditioning composition to dry, untreated substrate (fibre plus binder). Preferably, the amount of the conditioning composition ranges from about 5:1 to about 1:1, most preferably from about 3:1 to 1:1, by weight of the dry, untreated substrate.

According to one preferred embodiment of the invention, the dryer sheet substrate is coated by being passed over a rotogravure applicator roll. In its passage over this roll, the sheet is coated with a thin, uniform layer of molten fabric softening composition contained in a rectangular pan at a level of about 15g/square yard. Passage of the substrate over a cooling roll then solidifies the molten softening composition to solid. This type of applicator is used to obtain a uniform homogeneous coating across the sheet.

Following application of the liquefied composition, the articles are held at room temperature until the composition substantially solidifies. The resulting dry articles, prepared at the composition substrate ratios set forth above, remain flexible; the sheet articles are suitable for packaging in rolls. The sheet articles can optionally

be slitted or punched to provide a non-blocking aspect at any convenient time if desired during the manufacturing process.

The following Examples will more fully illustrate the embodiments of this invention. All parts, percentages and proportions referred to herein and in the appended claims are by weight of the composition unless otherwise indicated.

Examples 1-6 include organosilicones within the scope of the present invention having formulas A, B, C and D:

Formula A

 $\begin{array}{c} \text{si}(\text{CH}_3)_3 \\ \text{O} \\ \text{Si} \longrightarrow \text{osi}(\text{CH}_3)_3 \\ \text{(CH}_2)_3 \\ \text{N-[CH}_2 - \text{CH-(CH}_2)_5 - \text{CH}_3)_2 \\ \text{OH} \end{array}$

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Formula B

Formula C

Formula D

EXAMPLE 1

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The silicone of Formula C is a reaction product of the starting aminosilicone (where the nitrogen-containing branch chain is -(CH₂)₃-NH-(CH₂)₂-NH₂) and 1,2 epoxyoctadecane. The compound was prepared by placing the starting aminosilicone (61.16g), 1,2 epoxyoctadecane (38.84g) and 2-propanol (60.0g) in a reaction vessel and heating to 80°C for 24 hours. The reaction vessel consisted of a three neck round bottom flask containing a stirer, a reflux condenser and a thermometer. The 2-propanol was then stripped off with a N₂ sparge at 100°C as described in the Lin et al. application mentioned above.

Formula C silicone has %CH2 equal 56.62.

EXAMPLE 2

A 'T' structure modified alkylaminosilicone of Formula D, having %CH₂ equal 52.50 was prepared. In the starting aminoalkylsilicone, the nitrogen-containing branch chain is -(CH₂)₃-NH-(CH₂)₂NH₂. In the modified aminoalkylsilicone hydrogens on nitrogens were replaced with

OH

In the process, 34.7g of the starting aminoalkylsilicone, 34.4g 1,2-epoxydodecane and 17.4g 2-propanol were charged to the reaction vessel following the procedures of Example 1.

EXAMPLE 3

Effect of the %CH₂ content of various silicones as indicated in Table I on the compatibility with Adogen 442 (dihydrogenatedtallow dimethyl ammonium chloride from Sherex Corp.) was investigated by mixing the silicones with Adogen 442, following the SSCT procedure.

The results that were generated are summarised in Table I. Samples 6 and 7 were synthesised in Examples 1 and 2 respectively.

Table I

5	#	Silicone	%СН ₂	Compatible
	1.	DC 200 ¹	0	no
	2.	DC SSF ²	0	no
10	з.	Formula A	56.69	yes
	4.	Formula B	57.61	yes
	5.	Formula B, protonated	57.61	yes
15	6.	Formula C	56.62	yes
	7.	Formula D	52.50	yes

¹Linear polydimethylsiloxane, supplied by Dow Corning, viscosity = 1000cst

Silicones of samples 3 - 7 were mutually soluble with Adogen 442 at silicone concentration of 5% by weight of the mixture. However, silicone 1 and 2, which are not within the scope of the present invention, were not compatible with Adogen 442 at 5% or even at 25% of silicone.

Examples 4-6

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Contact Angle Measurements

Contact angle values reflect the spreading behaviour of a liquid on a solid surface. Discussion of the relationship between contact angle values and spreading is provided, for example, in Chapter 6 of "Introduction to Colloid and Surface Chemistry", Duncan J. Shaw, Butterworth, 1985. A contact angle of a liquid on solid surface is the angle between the tangent of the droplet and the surface. A smaller contact angle indicates better spreading on the surface. When it is desired to measure the contact angle on fabrics, there is an experimental problem of accurately measuring the true contact angle: due to the surface roughness of the fabric it is difficult to obtain an accurate baseline. Thus, the true contact angle measurements were obtained using cellulose paper.

Samples were prepared by mixing a fabric softener and a silicone above the melting point. A droplet of the melt liquid was applied to a piece of cellulose filter paper. After the droplet cooled and solidified, an initial contact angle was measured. The cellulose paper with the droplet was then piaced in a 70°C oven for 30 minutes in order for the equilibrium contact angle to be achieved. The paper was then removed from the oven and a final contact angle was measured.

The contact angle was measured using a contact angle goniometer (Rame'-Hart model 100). The cellulose with the drop of active was placed on the stage and viewed with a microscope. With the light source on, the drop appeared as a silhouette against a soft, green background. The drop/cellulose interface was aligned with the horizontal crosshair, and the contact angle was determined by rotating the read-out crosshair to tangency with the drop right profile. The contact angle value was then read directly on the graduated goniometer scale. This procedure was repeated to read the contact angle on the left side. Both sides should give the same reading otherwise the sample was not levelied correctly and the stage height should be readiusted.

55 Example 4

Effect of various silicones as indicated in Table II on the spreading of Adogen 442 was investigated. The true contact angle (initial and final) of the mixtures of silicones and Adogen 442 prepared in Example 3 was

²Aminosilicone supplied by Dow Corning, amine neutral equivalent = 2000, viscsity - 130cst.

measured on cellulose paper as described above. Additionally, spreading of the mixtures on cotton fabric was evaluated qualitatively, using a score of 1 to 4; 1 = best spreading, 2 = moderate spreading, 3 = droplet starting to wet the surface, 4 = no spreading, droplet beading up. Sample 1 contained only Adogen 442 without any silicone and was used as a control.

The results that were generated are summarised in Table II.

TABLE II

10	Sample	Silicone	Cotton	Celli	ılose
	No.			Initial	Final
15	1	none	4	110	112
15	2	DC200	4	110	147
	3	DCSSF	4	95	132
	4	Formula A	1	70	18
20	5	Formula B	2-3	72	59
	6	Formula B,			
		protonated	2	59	21
25	7	Formula C	2-3	86	57
	8	Formula D	1	47	60

Initial and final contact angles for samples 4-8 containing compatible silicones within the scope of the invention were lower than contact angles for samples 1-3.

Silicones of samples 4-8 were shown to form mutually soluble mixtures with Adogen 442 in Example 3. Samples 1-3 contained either no silicone or silicones which are not within the scope of the invention. The results established that, in mutually soluble mixtures of compatible silicones and fabric softener as taught by the present invention, compatible silicones improve the spreading of the fabric softener on a cellulose surface. Qualitative evaluation of spreading on cotton showed the same pattern of improved spreading when compatible silicones within the scope of the invention were used.

Example 5

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The concentration effect of various silicones as indicated in Table III on the spreading of Adogen 442 fabric softener was investigated by measuring the contact angle on a cellulose surface using the procedure described above.

TABLE_III

5	Sample No.	Silicone	Final Contact	Angle a	t % Silicone 6	of
10	1	DC200	147	147	147	
10	2	Formula B	118	55	_	
	3	Formula B,	,			
		protonated	1 48	20	20	
15	4	Formula D	42	-	55	
	5	Formula D,	,			
		protonated	98	78	5	

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This example demonstrates that in Samples 2-5 containing organosilicones within the scope of the invention as little as about 2% by weight of the mixture is needed to reduce the contact angle to improve the spreading on the surface.

Further increase in silicone concentration in Samples 2-5 further reduced the contact angle, indicating even better spreading on the surface.

Silicone of sample 1 which is not suitable for the present invention did not reduce the contact angle of the fabric softener regardless of the amount of the silicone used.

EXAMPLE 6

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Mixtures of various silicones as indicated in Table IV with nonionic fabric softeners, such as mineral oil were investigated. The spreading of the mixtures on cotton and polycotton fabrics was investigated by measuring the fabric area (centimetres²) per gram of mineral oil spread on the fabrics.

All sampples contained 5% by weight of the mixture of a silicone. The mineral oil used was Semtol 350 from Witco Corp.

TABLE IV

40	Sample No.	Silicone	Viscosity (cst)	Surface Tensio		ic Area Polycotton
45	1	none	105	32.0	303	371
	2	Formula B	295	22.9	227	224
	3	Formula D	182	22.2	326	552

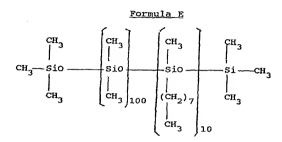
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Formula B silicone was only partially soluble in mineral oil, while Formula D silicone formed a mutually soluble mixture with mineral oil, demonstrating that the mutual solubility of the silicones and fabric softeners depend on the particular fabric softener as well as the %CH₂ of the silicone.

Silicones B and D both reduced the surface tension of mineral oil as observed in the absence of silicones in sample 1. However, fabric area coverage was increased only in sample 3 where a mutually soluble mixture was formed.

EXAMPLES 7-8

Examples 7-8 include organosilicones within the scope of the invention having formulas E, F and G.



Formula F

Formula G

EXAMPLE 7

The mutual solubility of organositicones with mixtures of fabric softening agents was investigated in the following formulations:

EP 0 459 822 A2

_	Formulation No.	Fabric Softening Component Mixture
5		
	r	10% Varisoft 475 ¹
10		10% Mineral Oil
	II	10% Adogen 442
		1% Myristic Acid
	III	11.7% Varisoft 445 ²
15		3.5% Stearic Acid
20	² Varisoft 445 = Methyl-1-hydr	n methyl sulfate
25	methyl sulfat	

The fabric softening mixtures of Formulations I, II and III above were heated and melted at approximately 80°C. Various silicones as indicated in Table V were added, with stirring, until the resulting mixture became hazy. At this point, the % silicone added was recorded as solubility of the silicone in the formulation. The results that were generated are summarised in Table V.

TABLE V

35	Formulation No.	PDMS ¹	Silicone Sol Silicone E	ubility (%) Silicone F
40				
	I	0.26	1.28	4.70
	II	0.34	0.69	3.10
4 5	III	0.39	1.69	15.58

¹PDMS = Polydimethylsiloxane, viscosity = 10,000 cst

Silicones E and F were significantly more soluble in Formulations I, II and III than PDMS.

Example 8

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Various silicones within the scope of the invention as indicated in Table VI were incorporated into liquid fabric conditioning compositions. Fabric softening agents and silicones were mixed together at 80°C (above the melting point) and then dispersed into water at 60°C-80°C to form liquid compositions containing composite particles of the fabric softening component and the silicone.

The resulting compositions are summarised in Table VI.

TABLE_VI

Ingredients	Sample A	В	С	D	E	F	G	
								
Adogen 442	7.3	7.3	-	-	- 1	.3.3	_	
Varisoft 475	-	-	10	10	10	_	-	
Varisoft 445	-	_	-	_	~	-	11.7	11
Neodol 23 ¹	0.94	0.94	_	-	-	_	_	
Siponic L7-90 ²	0.94	0.94	- "	-	_	-	_	
Mineral Oil	-	-	10	10	10	-	-	
Myristic acid		_	_	_	-	1.25	_	
Stearic acid		-	-	-	_	_	3.5	3
Silicone E	0.119	-	0.2	_	_	0.131	0.21	L3
Silicone G	-	0.1	-	0.2	_	_	_	
Silicone F	_		-	-	0.2	-	-	1
Water	90.7	90.7	79.8	79.8	79.8	85.3	84.6	5
82.9								

²Siponic L-7-90 = $C_{12}H_{25} - (OCH_2CH_2)_{12}OH$, from Alcolac.

Samples C, D, E, G and H were further tested for their softening properties. Terry cloths were prewashed with a solution of Neodol 25-9 (alcohol ethoxylate from Shell Corp.) and Na₂CO₃ to remove textile finishes on the surface, rinsed with the samples in a Tergometer and then line-dried. The cloth load was 20g per litre and the active concentration was 0.1g per litre of rinse liquid. The control was rinsed with only water. Using paired comparison, a panel of 20 judges assessed the softness of the treated cloth vs. control. All panelists preferred the treated cloths over the control in all tests.

EXAMPLES 9-11

The compatibility of various fabric softening agents with various silicones was determined by the SSCT. The entire concentration range up to 100% of the silicones was investigated. Samples that remained clear over the entire range of silicone concentration were labelled 'completely soluble'. For samples that became cloudy stability of the dispersions was ascertained and a and p compatibility values were determined by the SSCT.

The silicones that were investigated are listed in Table II. In the silicone formulas of Table II. $M = Me_3SIO_{0.5}$,

D = Me2Si-O,

and R' is as indicated in Table II.

Table II

5	Code	Formula	R*	%СН2
10	AA	Polydimethylsiloxane		
10		(= 1000 cst)	_	0
	BB	MD100D*5M	C8H17	14
	CC*	MD100D*5M	C ₁₈ H ₃₇	28
15	DD	MD300D*20M	C ₁₈ H ₃₇	28
	EE	MD100D*10M	C ₁₈ H ₃₇	43
	FF	MD95D*24M	C ₁₂ H ₂₅	57

*Code CC, Formula MD100D*5M is equivalent to Formula G of Example 8.

25 Example 9

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In this example, mixtures of the silicones listed in Table II with mineral oil were investigated using the SSCT. The mineral oil used was Fished Light Mineral oil. The results that were generated are summarised in Table III.

<u>Table III</u>

Compatibility with Mineral Oil

40	SILICONE	α COMPATIBILITY	β COMPATIBILITY	COMPATIBLE (YES/NO)
	AA	1	95	NO
	вв	4	80	NO
45	CC	COMPLETELY	SOLUBLE	YES
	EE	COMPLETELY	SOLUBLE	YES
	FF	COMPLETELY	SOLUBLE	YES

As determined by the SSCT, silicones CC, EE and FF having the structural requirements and %CH₂ recited by the present invention form compatible mixtures with mineral oil.

Example 10

In this example, mixtures of the silicones listed in Table II with various cationic quaternary fabric softening agents were investigated using the SSCT.

The results that were generated are summarised in Tables IV, V and VI.

TABLE IV

Compatibility with Varisoft 1371

SILICONE	α COMPATIBILITY	eta	COMPATIBLE (YES/NO)
22	2	97	NO
BB	2	98	NO
CC	2	96	ио
EE	7	93	YES
FF	7	90	YES

¹Varisoft 137 = di(hydrogenated)tallow dimethyl ammonium methylsulfate from Sherex.

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TABLE V

Compatibility with Varisoft 4451

SILICONE	α COMPATIBILITY	β COMPATIBILITY	COMPATIBLE (YES/NO)
AA	2	97	NO
EE	10	97	YES
FF	_	97	YES

¹Varisoft 445 = di(hydrogenated)tallow imidazolinium methylsulfate from Sherex.

TABLE V

Compatibility with Varisoft 1101

SILI	CONE	COMPATIBILITY	β COMPATIBILITY	(YES/NO)
AA		1	98	NO .
EE		5	90	YES
FF		5	90	YES

amidoethyl)

2-hydroxyethyl ammonium methylsulfate from Sherex

Example 11

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In this example, mixtures of the silicones listed in Table II with various nonionic fabric softening agents were investigated using the SSCT.

Results that were generated are summarised in Table VII, VIII, IX and X.

Table VII

Compatibility with Neodol 45-71

40	SILICONE	α COMPATIBILITY	β COMPATIBILITY	COMPATIBLE (YES/NO)
	AA	1	99	мо
45	вв	1	99	МО
	DD	2	99	мо
	FF	5	93	YES

1Neodol 45-7 = ethoxylated fatty alcohol from Shell.

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Table VIII

Compatibility with Adogen $345D^{1}$

SILICONE	α COMPATIBILITY	β COMPATIBI	COMPATIBLE LITY (YES/NO)	
A	2	60	NO	
В	COMPLETELY	SOLUBLE	YES	
D	COMPLETELY	SOLUBLE	YES	
E	COMPLETELY	SOLUBLE	YES	
F	COMPLETELY	SOLUBLE	YES	
1Adogen 34 Sherex.	45D = di(hydrogenate	ed)tallow	dimethyl amine	from

Table IX

Compatibility with PEG 600¹

35	SILICONE	compatibility	β COMPATIBILITY	COMPATIBLE (YES/NO)
	AA	2	. 99	мо
40	ВВ	2	98	NO
	DD	4	95	NO
	EE	4	95	YES
45	FF	4	95	YES

¹PEG 600 = Polyethylene Glycol

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Table X

Compatibility with isostearic acid

10	SILICONE	α COMPATIBILITY	$oldsymbol{eta}$ COMPATIBILITY	COMPATIBLE (YES/NO)
	AA	3	95	мо
	FF	3	96	YES

Examples 3-6 demonstrate that mutual compatibility between the fabric softening component and organosilicones may be easily determined by the SSCT and that the compatibility depends on the structure and %CH₂
content of the silicone as well as the particular fabric softening component employed in the mixture. Although
silicone C was highly compatible (mutually soluble) with mineral oil in Example 3 and with Adogen 345D in
Example 6, it was less compatible with Varisoft 137 of Example 4, i.e. a cloudy mixture was formed at 2% of
silicone. However, silicone C was more compatible with Varisoft 137 in Example 4 than polydimethylsioxane,
since β compatibility was lower for silicone C than for polydimethylsioxane. Results in Table VIII indicate that
amines have the highest degree compatibility with organosilicones, since silicone B, which has the %CH₂ content of 14% and is not within the scope of this invention is still compatible with di(hydrogenated)tallow dimethyl
amine. Silicones E and F, having a high %CH₂ content (43% and 57% respectively) were the most compatible
with all softeners tested.

30 EXAMPLE 12

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Two fabric softening sheets, A and B were prepared as follows:

The ingredients of a fabric conditioning composition as listed below were mixed in the melt. 500g of the prepared fabric conditioning mixture was placed in the pan of a two-roll coating machine and coated onto a spun-bonded polyester non-woven material. The fabric softening articles thus manufactured contained about 1.6g of solidified softening composition. The articles of manufacture were then placed into a tumble dryer machine which already contained 2.2 kg of prewashed clothing, including terry towelling softness monitors. The fabrics were then tumble dried with the fabric softening article until dry and the softening benefit was evaluated by a 20 member panel.

Fabric Conditioning formulation for sheet A:

- a. 10% of a silicone note suitable for use in the present invention (silicone B from Table II)
- b. 70% di(hydrogenated)tallow dimethyl ammonium methylsulfate
- c. 20% stearic acid.

Fabric Conditioning formulation for sheet B:

- a. 7% of a silicone within the scope of this invention (silicone FF from Table II)
- b. 70% di(hydrogenated tallow dimethyl ammonium methylsulfate
- c. 23% stearic acid.

Observations and results:

Sheet A - Due to the incompatible nature of the silicone, the silicone separated from the softening component during the coating process. The articles thus contained unknown amounts of the silicone.

Sheet B. The compatible silicone of the invention and the softening component formed a compatible mixture which remained homogeneous during the coating process as it was transferred to the substrate indicating that the substrate was uniformly and evenly coated.

A 20 member panel judged the towelling monitors for both sheet A and sheet B to have superior softness vs. towels prepared in an identical fashion but dried without softener.

This invention has been described with respect to certain preferred embodiments and various modifications

thereof will occur to persons skilled in the art in the light of the instant specification and are to be included within the spirit and purview of this application and the scope of the appended claims.

5 Claims

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- A liquid fabric conditioning composition comprising about 1% to about 60% of composite particles consisting of a compatible mixture consisting of:
 - a. about 1% to about 40% by weight of the conditioning composition of a fabric softening component comprising a cationic quaternary ammonium salt; and
 - b, about 0.1% to about 20% by weight of the conditioning composition of an organosilicone having a %CH₂ content of about 25% to about 90% and having at least one unit of Formula A:

$$R^{1} - \operatorname{Si} - O_{(3-m)/2}$$

- wherein m is a number from 0 to 2, R is a mono valent hydrocarbon radical and R^1 is selected from the group consisting of:
- i. a hydrocarbon radical having from 6 to 45 carbon atoms;
- ii. a unit of Formula A1

wherein a is a number of at least 1, b is a number from 0 to 10,

$$R^2$$
 is -CH-CH- R^3 , R^3 is
$$\begin{cases} & \downarrow \\ & \downarrow \\ & R^4 \text{ OH} \end{cases}$$

a hydrocarbon radical having from 4 to 40 carbon atoms and R^{+} is hydrogen or a hydrocarbon radical having from 1 to 40 carbon atoms; and iii. a unit of Formula A2

$$-R^7 - N - R^5$$

wherein R⁵ and R⁶ are independently selected from hydrogen or a hydrocarbon radical having fro 1 to 45 carbon atoms and at least one of R⁵ and R⁶ is a hydrocarbon radical having from 6 to 45 carbon atoms, R⁷ is

$$-R^8$$
 - CH - CH₂ - ,

where R8 is a divalent organic radical having from 1 to 12 carbon atoms.

- The composition of claim 1 wherein the %CH₂ content of said organosilicone is about 40% to about 90%.
 - The composition of claim 1 or claim 2 wherein an amount of said organosillcone is about 2% to about by weight of said composition.
- 4. The composition of any preceding claim wherein R¹ includes from 8 to 18 carbon atoms.
 - 5. The composition of any preceding claim wherein a is 3 and b is 1.
 - 6. The composition of any preceing claim wherein R3 includes from 8 to 18 carbon atoms.
 - 7. The composition of any preceding claim wherein R4 is hydrogen.
 - 8. The composition of any preceding claim wherein m is 1.
- The composition of any preceding claim wherein the composite particles consist of a mutually soluble mixture.
 - Method for softening fabrics comprising treating said fabrics in an aqueous bath with the fabric conditioning composition of claim 1.
 - 11. Discrete composite particles consisting of a compatible mixture consisting of:
 - a, at least about 1% of a fabric softening component comprising a cationic quaternary ammonium salt; and
 - b. an organosilicone having a %CH₂ content of about 25% to about 90% and having at least one unit of Formula A:

$$R^1 - Si^{-0}(3-m)/2$$

wherein m is a number from 0 to 2, R is a mono valent hydrocarbon radical and R^1 is selected from the group consisting of:

i. a hydrocarbon radical having from 6 to 45 carbon atoms;

ii. a unit of Formula A1

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wherein a is a number of at least 1, b is a number from 0 to 10, R2 is

R³ Is a hydrocarbon radical having from 4 to 40 carbon atoms and R⁴ is hydrogen or a hydrocarbon radical having from 1 to 40 carbon atoms; and iii. a unit of Formula A2

 $-R^7 - N - R^5$ $\begin{matrix} & & & \\ &$

wherein R⁵ and R⁶ are independently selected from hydrogen or a hydrogen radical having from 1 to 45 carbon atoms and at least one of R⁵ and R⁶ is a hydrocarbon radical having from 6 to 45 carbon atoms, R⁷ is

$$-R^8$$
 - CH - CH_2 - , I

where R8 is a divalent organic radical having from 1 to 12 carbon atoms.

- 12. An article for conditioning fabrics which provides for release of a fabric conditioning composition within an automatic laundry dryer at dryer operating temperatures comprising a flexible substrate and an amount effective to condition fabrics of a fabric conditioning composition, carried on said substrate, said composition comprising:
 - a. a fabric softening component selected from:
 - i, cationic quaternary ammonium saits:
 - ii. nonkonic fabric softeners selected from the group consisting of tertiary amines having at least one C_{8-30} alkyl chain, esters of polyhydric alcohols, fatty alcohols, ethoxylated fatty alcohols, alkylphenols, ethoxylated alkylphenols, ethoxylated monoglycerides, ethoxylated diglycerides, ethoxylated fatty amines, mineral oils, polyols, and mixtures thereof;
 - iii. carboxylic acids having at least 8 carbon atoms; and
 - iv. mixtures thereof; and

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b, an organosilicone having a % CH₂ content of about 25% to about 90% and having at least one unit of Formula A;

R¹ - Si-O_{(3-m)/2}

wherein m is a number from 0 to 2, R is a mono valent hydrocarbon radical and R^1 is selected from the group consisting of:

- i. a hydrocarbon radical having from 6 to 45 carbon atoms;
- ii. a unit of Formula A1

$$-(CH_2)_a - (N-CH_2CH_2)_b - N-R^2$$

wherein a is a number of at least 1, b is a number from 0 to 10, R2 is

 R^s is a hydrocarbon radical having from 4 to 40 carbon atoms and R^s is hydrogen or a hydrocarbon radical having from 1 to 40 carbon atoms; and iii. a unit of Formula A2

wherein R⁵ and R⁶ are independently selected from hydrogen or a hydrocarbon radical having from 1 to 45 carbon atoms and at least one of R⁵ and R⁶ is a hydrocarbon radical having from 6 to 45 carbon atoms, R⁷ is

$$-R^8$$
 - CH - CH_2 - ,

where R⁸ is a divalent organic radical having from 1 to 12 carbon atoms, wherein said fabric softening component and said organosilicone form a compatible mixture.

- 13. The article of claim 12 wherein the %CH2 content of said organosilicone is about 40% to about 90%.
- 14. The article of claim 12 or claim 13 wherein an amount of said organosilicone is about 0.1% to about 20% by weight of said composition.
 - 15. The article of claim 12 or claim 13 or claim 14 wherein an amount of said organosilicone is about 3% to about 20% by weight of said composition.
 - 16. The article of any of claims 12 to 15 wherein said flexible substrate is in a sheet configuration.
 - A method of conditioning laundry in a dryer comprising contacting said laundry with the fabric conditioning article of claims 12 to 16.

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